Connecting via Winsock to Dialog

Logging in to Dialog Trying 31060000009998...Open DIALOG INFORMATION SERVICES PLEASE LOGON: ****** ENTER PASSWORD: ****** Welcome to DIALOG Dialog level 05.26.00D Last logoff: 12aug09 09:23:50 Logon file405 12aug09 15:10:11 SYSTEM: HOME Cost is in DialUnits Menu System II: D2 version 1.8.0 term=ASCII *** DIALOG HOMEBASE(SM) Main Menu *** Information: 1. Announcements (new files, reloads, etc.) 2. Database, Rates, & Command Descriptions 3. Help in Choosing Databases for Your Topic 4. Customer Services (telephone assistance, training, seminars, etc.) 5. Product Descriptions Connections: 6. DIALOG(R) Document Delivery 7. Data Star(R) (c) 2003 Dialog, a Thomson business. All rights reserved. /H = Help/L = Logoff /NOMENU = Command Mode Enter an option number to view information or to connect to an online service. Enter a BEGIN command plus a file number to search a database (e.g., B1 for ERIC). 2 h 410 12aug09 15:10:11 User226352 Session D1162.1 \$0.00 0.267 DialUnits FileHomeBase \$0.00 Estimated cost FileHomeBase \$0.00 Estimated cost this search

```
$0.00 Estimated total session cost 0.267 DialUnits
File 410: The Chronolog 2009
       (c) 2009 Dialog. All rights.reserved.
     Set Items Description
      ---
? set hi ;set hi
HILIGHT set on as ''
HILIGHT set on as ''
? b biochem
       12aug09 15:10:18 User226352 Session D1162.2
                   0.117 DialUnits File410
           $0.00
    $0.00 Estimated cost File410
    $0.02 TELNET
    $0.02 Estimated cost this search
    $0.02 Estimated total session cost 0.384 DialUnits
SYSTEM:OS - DIALOG OneSearch
 File 5:Biosis Previews(R) 1926-2009/Aug W2
        (c) 2009 The Thomson Corporation
 File
       6:NTIS 1964-2009/Aug W4
        (c) 2009 NTIS, Intl Covraht All Rights Res
 File 24:CSA Life Sciences Abstracts 1966-2009/Aug
        (c) 2009 CSA.
 File 34:SciSearch(R) Cited Ref Sci 1990-2009/Aug W1
        (c) 2009 The Thomson Corp
 File 40:Enviroline(R) 1975-2008/May
         (c) 2008 Congressional Information Service
*File 40: This file is closed and will no longer update. For
similar data, please search File 76-Environmental Sciences.
 File 41:Pollution Abstracts 1966-2009/Aug
        (c) 2009 CSA.
 File 45:EMCare 2009/Aug W1
        (c) 2009 Elsevier B.V.
 File 50:CAB Abstracts 1972-2009/Aug W2
        (c) 2009 CAB International
 File 65: Inside Conferences 1993-2009/Aug 12
        (c) 2009 BLDSC all rts. reserv.
 File 71:ELSEVIER BIOBASE 1994-2009/Aug W2
         (c) 2009 Elsevier B.V.
*File 71: The file has been reloaded. Accession numbers
have changed.
 File 72:EMBASE 1993-2009/Aug 10
         (c) 2009 Elsevier B.V.
*File 72: EMBASE Classic (File 772) now open to all Dialog customers.
See HELP NEWS 772 for information.
 File 73:EMBASE 1974-2009/Aug 10
        (c) 2009 Elsevier B.V.
*File 73: EMBASE Classic available to all Dialog customers.
See HELP NEWS 772 for information.
 File 76:Environmental Sciences 1966-2009/Aug
```

```
(c) 2009 CSA.
 File 98:General Sci Abs 1984-2009/Aug
         (c) 2009 The HW Wilson Co.
 File 103:Energy SciTec 1974-2009/Jul B2
         (c) 2009 Contains copyrighted material
*File 103: For access restrictions see Help Restrict.
 File 136:BioEngineering Abstracts 1966-2007/Jan
         (c) 2007 CSA.
*File 136: This file is closed.
 File 143:Biol. & Agric. Index 1983-2009/Jul
         (c) 2009 The HW Wilson Co
 File 144:Pascal 1973-2009/Aug W2
         (c) 2009 INIST/CNRS
 File 154:MEDLINE(R) 1990-2009/Aug 11
         (c) format only 2009 Dialog
 File 155:MEDLINE(R) 1950-2009/Aug 11
         (c) format only 2009 Dialog
 File 156:ToxFile 1965-2009/Aug W2
         (c) format only 2009 Dialog
 File 162:Global Health 1983-2009/Aug W2
         (c) 2009 CAB International
 File 172:EMBASE Alert 2009/Aug 11
         (c) 2009 Elsevier B.V.
 File 305: Analytical Abstracts 1980-2009/Jun W4
         (c) 2009 Roval Soc Chemistry
*File 305: Alert feature enhanced for multiple files, duplicate
removal, customized scheduling. See HELP ALERT.
 File 369: New Scientist 1994-2009/Aug W1
         (c) 2009 Reed Business Information Ltd.
 File 370:Science 1996-1999/Jul W3
         (c) 1999 AAAS
*File 370: This file is closed (no updates). Use File 47 for more
current
information.
 File 393:Beilstein Database - Abstracts 2008/02
         (c) 2008 Beilstein GmbH
 File 399:CA SEARCH(R) 1967-2009/UD=15107
         (c) 2009 American Chemical Society
*File 399: Use is subject to the terms of your user/customer
agreement.
IPCR/8 classification codes now searchable as IC=. See HELP NEWSIPCR.
 File 434:SciSearch(R) Cited Ref Sci 1974-1989/Dec
         (c) 2006 The Thomson Corp
     Set Items Description
      --- -----
? s (ysbC or orotate or orotic) (5n) (transpor?)
              12 YSBC
           9263 OROTATE
           19537 OROTIC
         7897667 TRANSPOR?
             203 (YSBC OR OROTATE OR OROTIC) (5N) (TRANSPOR?)
```

```
? rd s1
>>>Duplicate detection is not supported for File 393.
>>>Records from unsupported files will be retained in the RD set.
     S2
              97 RD S1 (unique items)
? s s2 not py>2006
              97
                 S2
        20271259 PY>2006
              92 S2 NOT PY>2006
? s s3 and (gene or nucleic or clone or polynucleic or DNA)
Processing
Processed 20 of 29 files ...
Completed processing all files
              92 53
        10933603 GENE
         1320192 NUCLEIC
          557564 CLONE
             664 POLYNUCLEIC
         8185040 DNA
     S4
              2.6
                 S3 AND (GENE OR NUCLEIC OR CLONE OR POLYNUCLEIC OR
DNA)
? t. s4/7/al
>>>'AL' not allowed as item list
? t s4/7/all
>>>Format 7 is not valid in file 143
           (Item 1 from file: 5)
DIALOG(R)File 5:Biosis Previews(R)
(c) 2009 The Thomson Corporation. All rts. reserv.
19125140 BIOSIS NO.: 200600470535
Mutations and rearrangements in the genome of Sulfolobus solfataticus
AUTHOR: Redder Peter (Reprint); Garrett Roger A
AUTHOR ADDRESS: Inst Pasteur, Unite Biol Mol Gene Extremophiles, 25
Rue Dr
 Roux, F-75724 Paris 15, France**France
AUTHOR E-MAIL ADDRESS: predder@pasteur.fr
JOURNAL: Journal of Bacteriology 188 (12): p4198-4206 JUN 2006 2006
TSSN: 0021-9193
DOCUMENT TYPE: Article
RECORD TYPE: Abstract
LANGUAGE: English
ABSTRACT: The genome of Sulfolobus solfataricus P2 carries a larger
number
 of transposable elements than any other sequenced genome from an
```

rearrangement and change. In order to gain more insight into the natures $% \left(1\right) =\left(1\right) +\left(1\right) +\left$

or bacterium and, as a consequence, may be particularly susceptible

archaeon

and frequencies of different types of mutation and possible rearrangements that can occur in the genome, the pyrEF locus was

rearrangements that can occur in the genome, the pyrEF locus was examined

for mutations that were isolated after selection with 5-fluoroorotic acid. About two-thirds of the 130 mutations resulted from

insertions of

mobile elements, including insertion sequence (IS) elements and a single $\,$

nonautonomous mobile element, SM2. For each of these, the element

identified and shown to be present at its original genomic position, consistent with a progressive increase in the copy numbers of the mobile

elements. In addition, several base pair substitutions, as well as small

deletions, insertions, and a duplication, were observed, and about one-fifth of the mutations occurred elsewhere in the genome, possibly in

an orotate transporter gene. One mutant exhibited a

 $5-{
m kb}$ genomic rearrangement at the pyrEF locus involving a two-step TS

element-dependent reaction, and its boundaries were defined using a specially developed "in vitro library" strategy. Moreover, while searching for the donor mobile elements, evidence was found for two major

changes that had occurred in the genome of strain P2, one constituting a

single deletion of about 4% of the total genome (124 kb), while the other $\,$

involved the inversion of a 25-kb region. Both were bordered by IS elements and were inferred to have arisen through recombination events

The results underline the caution required in working

experimentally with

an organism such as S. solfataricus with a continually changing genome. $% \left\{ \left(\frac{1}{2}\right) \right\} =\left\{ \left(\frac{1}$

4/7/2 (Item 2 from file: 5)

DIALOG(R)File 5:Biosis Previews(R)

(c) 2009 The Thomson Corporation. All rts. reserv.

13736133 BIOSIS NO.: 199799370193

Utilization of orotate as a pyrimidine source by Salmonella typhimurium and

Escherichia coli requires the dicarboxylate transport protein encoded by

dctA

AUTHOR: Baker Kristian E; Ditullio Katrina P; Neuhard Jan; Kelln Rod A (Reprint)

AUTHOR ADDRESS: Dep. Chem., Univ. Regina, Regina, SK S4S 0A2, Canada**
Canada

JOURNAL: Journal of Bacteriology 178 (24): p7099-7105 1996 1996 ISSN: 0021-9193

DOCUMENT TYPE: Article RECORD TYPE: Abstract

LANGUAGE: English

 $\label{eq:abstract: Mutants deficient in orotate utilization (initially termed out \\$

mutants) were isolated by selection for resistance to 5-fluoroorotate

(FOA), and the mutations of 12 independently obtained isolates were found

to map at 79 to 80 min on the Salmonella typhimurium chromosome. A gene complementing the mutations was cloned and sequenced and found to possess extensive sequence identity to characterized genes for C4-dicarboxylate transport (dctA) in Rhizobium species and to the sequence inferred to be the dctA gene of Escherichia coli. The mutants were unable to utilize succinate, malate, or fumarate as sole

carbon source, an expected phenotype of dctA mutants, and introduction of

the cloned DNA resulted in restoration of both C4-dicarboxylate and orotate utilization. Further, succinate was found to compete with orotate

for entry into the cell. The S. typhimurium dctA gene encodes a highly hydrophobic polypeptide of 45.4 kDa, and the polypeptide was found

to be enriched in the membrane fraction of minicells harboring a $\mbox{dct} A +$

plasmid. The DNA immediately upstream of the deduced -35 region contains a putative cyclic AMP-cyclic AMP receptor protein complex binding site, thus affording an explanation for the more effective utilization of orotate with glycerol than with glucose as carbon source.

The E. coli dctA gene was cloned from a lambda vector and shown to complement C4-dicarboxylate and orotate utilization in FOA-resistant mutants of both E. coli and S. typhimurium. The accumulated results demonstrate that the dctA gene product, in addition to transporting C4-dicarboxylates, mediates the transport of orotate, a cyclic monocarboxylate.

4/7/3 (Item 3 from file: 5)

DIALOG(R)File 5:Biosis Previews(R)

(c) 2009 The Thomson Corporation. All rts. reserv.

07201654 BIOSIS NO.: 198477033565

DIFFERENT RATES OF SYNTHESIS AND TURNOVER OF RIBOSOMAL RNA IN RAT BRAIN AND

LIVER

AUTHOR: STOYKOVA A S (Reprint); DUDOV K P; DABEVA M D; HADJIOLOV A A AUTHOR ADDRESS: INST MOLECULAR BIOLOGY, BULGARIAN ACAD SCI, 1113 SOFIA,

```
BULG**BULGARTA
JOURNAL: Journal of Neurochemistry 41 (4): p942-949 1983
TSSN: 0022-3042
DOCUMENT TYPE: Article
RECORD TYPE: Abstract
LANGUAGE: ENGLISH
ABSTRACT: The kinetics of in vivo labeling of cellular free UMP and
 nucleolar, nucleoplasmic, and cytoplasmic, rRNA [ribosomal RNA] with
  [14C]orotate in rat brain and liver were investigated. Evaluation
of the
 experimental data shows the following result: The rate of nucleolar
 precursors of ribosomal RNA (pre-rRNA) synthesis and the deduced
rate of
 ribosome formation in brain are about 5-fold lower than in liver and
  correspond to 220-260 ribosomes/min/nucleus. The lower rate of in
vivo
  pre-rRNA synthesis is correlated with a lower activity of RNA
polymerase
  I in isolated brain nuclei. The half-lives of nucleolar rRNA in
brain and
 liver are 210 and 60 min, respectively, thus showing a slower rate
 processing of pre-rRNA in brain nucleoli. The nucleo-cytoplasmic
 transport of ribosomes in brain is also markedly slower than in
 reflects the lower rates of synthesis and processing of pre-rRNA.
  Cytoplasmic ribosomes in brain and liver turn-over with half-lives
of
  about 6 and 4 days, respectively. The markedly lower rate of
ribosome
 biogenesis in brain is specified mainly at the level of
transcription of
 rRNA genes.
 4/7/4
           (Item 4 from file: 5)
DIALOG(R)File 5:Biosis Previews(R)
(c) 2009 The Thomson Corporation. All rts. reserv.
          BIOSIS NO.: 197967041713
PYRIMIDINE NUCLEOTIDE BIOSYNTHESIS A STUDY OF NORMAL AND PURINE ENZYME
 DEFICIENT CELLS
AUTHOR: FOX I H (Reprint); BURK L; PLANET G; GOREN M; KAMINSKA J
AUTHOR ADDRESS: UNIV MICH MED CENT, ANN ARBOR, MICH 48109, USA**USA
```

ABSTRACT: To evaluate the importance of altered pyrimidine synthesis in the $% \left(1\right) =\left(1\right) +\left(1\right)$

JOURNAL: Journal of Biological Chemistry 253 (19): p6794-6800 1978

ISSN: 0021-9258
DOCUMENT TYPE: Article
RECORD TYPE: Abstract
LANGUAGE: ENGLISH

relationship between immune dysfunction and the deficiencies of adenosine $% \left(1\right) =\left(1\right) \left(1\right) +\left(1\right) \left(1\right) \left(1\right) +\left(1\right) \left(1\right$

deaminase or purine nucleoside phosphorylase, this pathway was measured

in human erythrocytes or fibroblasts in vitro or in cultured diploid fibroblasts during growth. The effect of purine nucleosides on the conversion of orotic acid to UMP in intact erythrocytes or fibroblasts

was assayed by the release of CO2 from orotic acid. Adenosine caused $50 \, \$$

inhibition (I0.5) of CO2 production at 80 μM in erythrocytes and at 270 μM in fibroblasts. Quantitatively similar changes occurred in intracellular concentrations of PP-ribose-P. Studies of the mechanism for

this inhibition in erythrocytes suggest a regulatory role for PP-ribose-P

concentrations. Increases or decreases of erthrocyte PP-ribose-P concentrations were accompanied by similar changes in CO2 release from

orotic acid, suggesting that PP-ribose-P is rate-limiting for orotate $% \left(1\right) =\left(1\right) +\left(1$

phosphoribosyltransferase. Adenosine did not inhibit orotic acid transport into erythrocytes and did not directly inhibit orotate phosphoribosyltransferase or orotidylic decarboxylase. In cultured diploid fibroblasts, adenosine (50 or 100 μM) causes 73 or 76% inhibition, respectively, of the ratio of orotic acid to uridine incorporation into nucleic acid. There is no decrease of this ratio in cells deficient in purine nucleoside phosphorylase, adenosine deaminase or hypoxanthine-quanine phosphoribosyltransferase. Abparently

adenosine blocks pyrimidine biosynthesis at orotate phosphoribosyltransferase in erythrocytes and fibroblasts. The hypothesis

that a block of pyrimidine synthesis is the basis for the immune disorder

in patients with deficiencies of purine nucleoside phosphorylase or adenosine deaminase was not supported.

4/7/5 (Item 5 from file: 5)
DIALOG(R)File 5:Biosis Previews(R)
(c) 2009 The Thomson Corporation. All rts. reserv.

05618845 BIOSIS NO.: 197967007840

EFFECT OF HEPATIC DE NERVATION ON RNA LEVELS AND CARBON-14 OROTIC-ACID INCORPORATION INTO HEPATIC NUCLEAR RNA

AUTHOR: OPANASYUK N D (Reprint); MASYUK A I; BEZDROBNYI YU V AUTHOR ADDRESS: DIV PATHOL PHYSIOL, KIEV MED INST, KIEV, USSR*USSR JOURNAL: Fiziolohichnyi Zhurnal (Kiev) 23 (5): p683-685 1977 ISSN: 0015-3311

DOCUMENT TYPE: Article RECORD TYPE: Abstract LANGUAGE: UKRAINIAN

ABSTRACT: RNA level increased to 15% above control in Wistar rats on the

9th day after ligation of the plexus near the porta hepatis;

not affect DNA levels. The RNA/DNA ratio increased 1.34-fold. After 21 days, RNA levels dropped to preligation levels; the decrease in

DNA levels at this time was related to dystrophic changes in the liver. The RNA/DNA ration remained above control. The rate of 14C-orotic acid incorporation into the thermophenol fraction of nuclear

RNA was studied to determine causes of RNA increase on the 9th day. $14C\mbox{-}orotic$ acid incorporation into 40S RNA increased 1.8-fold while no

changes were noted in incorporation into 65S RNA. Changes in $14C\text{-}\mathrm{orotic}$

acid incorporation was related to RNA synthesis and orotic acid transport across the cell membrane. Increases in RNA levels after ligation are due to increased synthesis of ribosomal RNA and activation

of protein synthesis mechanisms in the cell. The nervous system regulates, therefore, hepatic metabolism through the hepatocyte genetic

system.

4/7/6 (Item 1 from file: 24)
DIALOG(R)File 24:CSA Life Sciences Abstracts
(c) 2009 CSA. All rts. reserv.

0002902845 IP ACCESSION NO: 6947339 Mutations and Rearrangements in the Genome of Sulfolobus solfataricus P2

Redder, Peter; Garrett, Roger A Danish Archaea Centre, Institute for Molecular Biology and Physiology, Copenhagen University, Soelvgade 83H, DK-1307 Copenhagen K, Denmark

Journal of Bacteriology, v 188, n 12, p 4198-4206, June 2006 PUBLICATION DATE: 2006

PUBLISHER: American Society for Microbiology, 1752 N Street N.W. Washington, DC 20036 USA, [URL:http://www.asm.org/]

DOCUMENT TYPE: Journal Article RECORD TYPE: Abstract LANGUAGE: English SUMMARY LANGUAGE: English ISSN: 0021-9193 ELECTRONIC ISSN: 1098-5530 FILE SEGMENT: Genetics Abstracts; Bacteriology Abstracts (Microbiology B)

ABSTRACT:

The genome of Sulfolobus solfataricus P2 carries a larger number of transposable elements than any other sequenced genome from an archaeon or

bacterium and, as a consequence, may be particularly susceptible to rearrangement and change. In order to gain more insight into the natures

and frequencies of different types of mutation and possible rearrangements

that can occur in the genome, the pyrEF locus was examined for mutations $% \left(1\right) =\left(1\right) +\left(1\right) +\left$

that were isolated after selection with 5-fluoroorotic acid. About two-thirds of the 130 mutations resulted from insertions of mobile elements, including insertion sequence (IS) elements and a single nonautonomous mobile element, SM2. For each of these, the element was identified and shown to be present at its original genomic position, consistent with a progressive increase in the copy numbers of the mobile

elements. In addition, several base pair substitutions, as well as small

deletions, insertions, and a duplication, were observed, and about one-fifth of the mutations occurred elsewhere in the genome, possibly in an

orotate transporter gene. One mutant exhibited a 5-kb genomic rearrangement at the pyrEF locus involving a two-step IS element-dependent reaction, and its boundaries were defined using a specially developed "in vitro library" strategy. Moreover, while searching

for the donor mobile elements, evidence was found for two major changes

that had occurred in the genome of strain P2, one constituting a single

deletion of about 4% of the total genome (124 kb), while the other involved the inversion of a 25-kb region. Both were bordered by IS elements

and were inferred to have arisen through recombination events. The results

underline

the caution required in working experimentally with an organism such as S.

solfataricus with a continually changing genome.

4/7/7 (Item 1 from file: 50)
DIALOG(R)File 50:CAB Abstracts
(c) 2009 CAB International. All rts. reserv.

0005541318 CAB Accession Number: 19851465564

Effects of long-term maternal protein restriction on liver DNA, RNA and protein metabolism and subcellular distribution in developing rat.

Lewis, C. G.; Cheng, M.; Winick, M. Inst. Human Nutrition, Columbia University, College of Physicians and

Surgeon, New York, NY 10032, USA.

Nutrition Reports International volume 30 (1): p.199-211 Publication Year: 1984

Language: English

Record Type: Abstract

Document Type: Journal article

Pregnant rats were fed on a 25% casein diet (control) or switched to a $\,$

 $6\ensuremath{\,\%}$ casein diet at the beginning of the final third of pregnancy and to a

10% casein diet after giving birth (malnourished). All rats were fed $\,$

freely and at birth all litters were reduced to 8. From 13 to 21 days of $\,$

age, total liver DNA (cell number) and the incorporation of [sup 3H]thymidine into DNA was significantly retarded in malnourished young. Total liver RNA and protein content were retarded in malnourished

young but cellular RNA and protein content and subcellular distribution

were not different from control values. There was no difference in

incorporation of label from [SUP 14 C]orotate into RNA, transport of RNA from nucleus to cytoplasm, subcellular fractional distribution or RNA half-life when compared to control values. It it

suggested that the 3 weeks of perinatal malnutrition represent long-term $\,$

nutritional deprivation to the rapidly growing young and that the liver $% \left(1\right) =\left(1\right) +\left(1\right) +\left($

has adapted to protein malnutrition by reducing total organ cell number to $% \left(1\right) =\left(1\right) +\left(1\right)$

maintain functional capacity and integrity of existing cells.

4/7/8 (Item 1 from file: 73)
DIALOG(R)File 73:EMBASE
(c) 2009 Elsevier B.V. All rts. reserv.

0071652423 EMBASE No: 1980095172

Biochemistry of Plasmodium (malarial parasites)

Sherman I.W.

Dept. Biol., University California, Riverside, Calif. 92521, United States:

CORRESP. AUTHOR/AFFIL: Dept. Biol., University California, Riverside, Calif.

92521, United States

```
Microbiological Reviews (MICROBIOL. REV.) (United States)
December 1,
1979, 43/4 (453-495)
CODEN: MBRED ISSN: 0146-0749
DOCUMENT TYPE: Journal; Review RECORD TYPE: Abstract
LANGUAGE: English
```

Intraerythrocytically, bird and mammalian malarias appear to derive energy by metabolizing glucose to lactic acid via a conventional pathway of

anaerobic glycolysis. If supplied with oxygen, avian parasites may oxidize

- a portion of the pyruvate to CO SUB 2 and water by means of the citric acid
- cycle, whereas rodent and primate malarias, which lack a functional citric
- acid cycle, are unable to do so and have lactate as their primary end product. Erythrocyte-free P. gallinaceum cells produce appreciable quantities of acetate from glucose and pyruvate. Since plasmodia are unable
- to synthesize CoA de novo, it is possible that in P. gallinaceum acetate
- formation is due to a lack of host-supplied CoA. In P.knowlesi it may be
- that volatile acids are formed by a pyruvate clastic reaction, but the enzymes involved (if they exist) have not been looked for. However, in view
- of the fragile nature of free parasites, acetate and formate production may
- reflect deranged metabolism due to in situ leakiness of plasmodia or may be
- a consequence of insult during the isolation procedure. There is no evidence for a pentose phosphate shunt in malarial parasites since the first enzyme in the pathway (GGPDH) is absent. Indeed, the only enzyme in
- this pathway that has been identified consistently is 6-phosphogluconate
- dehydrogenase. Lacking a pentose shunt, the parasites must have other means
- for obtaining ribose and reducing NADP. It has been suggested that action
- by phosphorylases supplies the pentoses and that glutamic dehydrogenase
- provides for the reduction of NADP. Evidence for an energy-yielding electron transport chain is at best circumstantial. In all of the malarias
- studied, the only enzyme found to be associated with this system is cytochrome oxidase. It is conceivable that in the acristate rodent malarias
- and in P. knowlesi, and perhaps even in those malarias having cristate mitochondria (avians and P. falciparum), cytochrome oxidase is involved in

the de novo pyrimidine biosynthetic pathway and not in energy-yielding reactions. Malarial parasites are incapable of de novo purine biosynthesis;

however, pyrimidines are synthesized de novo. Exogenously supplied purines

and orotic acid are transported and incorporated by infected erythrocytes and plasmodia, whereas pyrimidines (uracil and thymidine) are

 $\ensuremath{\operatorname{not}}.$ There is evidence to support the contention that hypoxanthine is the

preferred purine of the parasites in vivo and that it is derived from the

catabolism of erythrocytic ATF. Plasmodia have a distinctive DNA and rRNA base composition. Malarial parasite ribosomes are not provided for by

host cell ribosomal subparticles, and the mechanism of protein synthesis by

the parasites is typically eucaryotic. The capacity of the parasites for \mbox{de}

novo amino acid biosynthesis is limited, and it appears that host cell hemoglobin provides most of the amino acids. For some species, isoleucine

and methionine must be supplied exogenously for good plasmodial growth. The $\ensuremath{\,}^{}$

degradation of erythrocyte hemoglobin by parasite proteases leaves a golden

brown-black residue called hemozoin (malarial pigment). Hemozoin consists $% \left(1\right) =\left(1\right) \left(1\right) +\left(1\right) \left(1\right) \left(1\right) +\left(1\right) \left(1\right$

of insoluble monomers and dimers of hematin, methemoglobin, and ferriprotoporphyrin coupled to plasmodial protein. The functional significance of hemozoin is not completely understood. The only well-characterized plasmodial protein is the HRP of P. lophurae. It is possible that HRP is localized in the polar organelles of the merozoites

and is involved in the process of invasion. Information regarding the vitamin requirements of malarial parasites is scanty. Plasmodia are incapable of synthesizing CoA from pantothenate and rely on the host cell

for this cofactor; this may be one reason for their being obligate intracellular parasites. By contrast, Plasmodium can synthesize folate from pABA.

4/7/9 (Item 2 from file: 73)
DIALOG(R)File 73:EMBASE
(c) 2009 Elsevier B.V. All rts. reserv.

0070373498 EMBASE No: 1975157339

Incorporation of precursors and inhibitors of nucleic acid synthesis into hepatomas and liver of the rat
Lea M.A.; Bullock J.; Khalil F.L.; Morris H.P.

Dept. Biochem., New Jersey Med. Sch., Newark, N.J. 07103, United States:

CORRESP. AUTHOR/AFFIL: Dept. Biochem., New Jersey Med. Sch., Newark, N.J.

07103, United States

Cancer Research (CANCER RES.) December 1, 1974, 34/12 (3414-3420) CODEN: CNREA $\,$ ISSN: 0008-5472

DOCUMENT TYPE: Journal; Article RECORD TYPE: Abstract LANGUAGE: English

In order to evaluate the entry of nucleic acid precursors into transplanted hepatomas and to examine the relationship to growth rate, the

incorporation of isotope label was measured in tissue fractions. The low $% \left(1\right) =\left(1\right) +\left(1\right) +\left$

rate of incorporation of orotate into $\ensuremath{\mathtt{RNA}}$ of hepatomas in comparison with

liver was confirmed and found to occur even in the most slowly growing tumors. A similar pattern was observed for the incorporation of orotate $\,$

into the acid soluble fraction. Under appropriate conditions, all the hepatomas examined were able to achieve an orotate incorporation greater

than that in blood and several other tissues. Similar data were obtained 60

min after either i.p. or s.c. injections. Studies with several other molecules including dihydroorotate, uracil, uridine, thymidine, inorganic

phosphate, 5 fluorouracil, and hycanthone did not show such pronounced changes in hepatomas but did suggest that uptake of these compounds is less

in more rapidly growing liver tumors than in the slowly growing tumors.

From the unequal incorporation of different molecules into a given tumor, $% \left(1\right) =\left(1\right) +\left(1\right) +$

from temporal studies of uptake, and from a comparative uptake study of

s.c. and intrahepatic tumors, it was concluded that the vascular supply was

not the sole determinant for the relative uptake of orotate in different

tumor tissues. The data suggested that a transport mechanism for orotate may be impaired in hepatic neoplasia. As the regenerating liver did not show this transition, it does not appear to be an essential

feature of cellular proliferation.

4/7/10 (Item 3 from file: 73) DIALOG(R)File 73:EMBASE (c) 2009 Elsevier B.V. All rts. reserv. 0070179484 EMBASE No: 1974181078

Biochemical effects of miconazole on fungi I. Effects on the uptake and/or utilization of purines, pyrimidines, nucleosides, amino acids and

glucose by Candida albicans

Van Den Bossche H.

Dept. Comp. Biochem., Res. Laboratory, Janssen Pharmaceut., Beerse, Belgium:

CORRESP. AUTHOR/AFFIL: Dept. Comp. Biochem., Res. Laboratory, Janssen

Pharmaceut., Beerse, Belgium

Biochemical Pharmacology (BIOCHEM. PHARMACOL.) January 1, 1974, 23/4

(887-899)

CODEN: BCPCA ISSN: 0006-2952 DOI: 10.1016/0006-2952(74)90220-2

DOCUMENT TYPE: Journal; Article RECORD TYPE: Abstract

LANGUAGE: English

The antifungal and antibacterial drug miconazole has been shown to inhibit, at concentrations lower than those affecting growth, the transport

of adenine, guanine and lllw2=hypoxanthine by Candida albicans in suspension culture. The decrease in the incorporation of purines into nucleic acids seems to be the consequence of an inhibitory effect on their uptake into the cells. When the purines were replaced by adenosine.

deoxyadenosine and guanosine, miconazole increased the uptake and incorporation of the radioactivity derived from the nucleosides into macromolecules. The data suggest that the drug induced increase of nucleoside incorporation into nucleic acids is secondary to enhanced nucleoside transport. Miconazole also slightly affected the uptake of orotic acid. The transport of glucose, glycine and leucine was not affected by miconazole whereas in some way the drug affected glutamine

uptake. Studies on the distribution of miconazole and/or its metabolites in

the Candida cell indicate that in log phase cells most of the radioactivity

was found in the fraction containing cell walls and plasmalemma. In stationary phase cells the highest radioactivity was found in the fraction

which contained the microsomes. Although more information will be needed,

the data presented indicate that at low concentrations, miconazole acts

primarily on the yeast cell membranes (cell wall and plasmalemma) resulting

in a selective inhibition of the uptake of precursors of RNA and DNA (purines) and mucopolysaccharide (glutamine). Higher doses and longer

incubation periods also alter the activities of microsomal membranes.

Title: Incorporation of labeled ribonucleic acid precursors into

(Item 1 from file: 103)

INS-86-020998; EDB-86-121620

(c) 2009 Contains copyrighted material. All rts. reserv.

DIALOG(R)File 103:Energy SciTec

4/7/11

01797832

```
maternal
    and fetal rat tissues during pregnancy
Author(s): Dorko, M.E.; Hayashi, T.T.
Affiliation: University of Pittsburgh School of Medicine, PA
Source: Am. J. Obstet. Gynecol. (United States) v 4. Coden: AJOGA
Publication Date: Apr 1986
p 801-805
Language: English
Abstract: Tritium-labeled ribonucleic acid precursors, including
cytidine,
   uridine, and orotic acid, were injected into rats with dated
   pregnancies (14 to 21 days) and virgin rats. The acid-insoluble
counts
    indicating incorporation into fetal and placental tissues showed
that
   the highest incorporation occurred with cytidine, particularly
earlier
    in pregnancy. In contrast, uridine demonstrated a minor degree of
    incorporation but displayed facile and enhanced transplacental
passage
   with duration of pregnancy as represented by acid-soluble counts.
   Orotic acid was minimally used by both fetal and placental
tissues. The
    incorporation of labeled precursors into maternal liver, heart,
and
   kidney demonstrated varying responses during the course of
pregnancy.
 4/7/12
           (Item 1 from file: 154)
DIALOG(R) File 154: MEDLINE (R)
(c) format only 2009 Dialog. All rts. reserv.
17528180 PMTD: 16968882
 A fluoroorotic acid-resistant mutant of Arabidopsis defective
in the
uptake of uracil.
 Mourad George S; Snook Bryan M; Prabhakar Joshua T; Mansfield
Tvler A:
Schultes Neil P
 Department of Biology, Indiana University-Purdue University Fort
Wayne
(IPFW).
        2101 East Coliseum Blvd, Fort Wayne, IN 46805-1499,
USA.
```

```
mourad@ipfw.edu
 Journal of experimental botany (England) 2006, 57 (14)
p3563-73,
ISSN 0022-0957--Print Journal Code: 9882906
 Publishing Model Print-Electronic
 Document type: Journal Article; Research Support, Non-U.S.
Research Support, U.S. Gov't, Non-P.H.S.
 Languages: ENGLISH
 Main Citation Owner: NLM
 Record type: MEDLINE; Completed
 A fluoroorotic acid (FOA)-resistant mutant of Arabidopsis
thaliana was
isolated by screening M2 populations of ethyl methane
sulphonate
(EMS)-mutagenized Columbia seed. FOA resistance was due to a
nuclear
recessive gene, for1-1, which locates to a 519 kb region in
chromosome 5. Assays of key regulatory enzymes in de novo
pyrimidine
synthesis
           (uridine monophosphate synthase) and salvage
biochemistry
(thymidine kinase) confirmed that FOA resistance in for1-1/for1-1
plants
    not due to altered enzymatic activities. Uptake studies
was
usina
radiolabelled purines, pyrimidines, and [14C]FOA reveal that
for1-1/for1-1
plants were specifically defective in the uptake of uracil or
uracil-like
bases. To confirm such specificity, genetic crosses show that
FOR1 is a
distinct
         locus from FUR1 which encodes a deoxyuridine
nucleoside
transporter. In addition, for1-1/for1-1 plants were restored
to FOA
sensitivity by transformation with the Escherichia coli uracil
transporter
gene uraA driven by the cauliflower mosaic virus (CaMV) 35S promoter.
Molecular mapping studies reveal that FOR1 does not correspond
to loci
belonging to any of the six known nucleobase transporter
families
identified in the Arabidopsis genome. Moreover, FOR1 does not
appear to
regulate the transcript levels of either uracil
transporter-encoding loci
At2q03590 or At2q03530. The above results strongly suggest that the
mutant allele affects a transport mechanism that is specific for the
uptake
of uracil.
```

Record Date Created: 20061114
Record Date Completed: 20070227
Date of Electronic Publication: 20060912

4/7/13 (Item 2 from file: 154)
DIALOG(R)File 154:MEDLINE(R)

(c) format only 2009 Dialog. All rts. reserv.

16589069 PMID: 15932997 Record Identifier: PMC1151845 sacB-5-Fluoroorotic acid-pyrE-based bidirectional selection for

integration of unmarked alleles into the chromosome of Rhodobacter capsulatus.

Yano Takahiro; Sanders Carsten; Catalano John; Daldal Fevzi Johnson Research Foundation, Department of Biochemistry and Biophysics,

School of Medicine, University of Pennsylvania, Philadelphia, Pennsylvania

19104, USA. vano@mail.med.upenn.edu

Applied and environmental microbiology (United States) $\,$ Jun 2005, 71 $\,$

(6) p3014-24, ISSN 0099-2240--Print Journal Code: 7605801 Contract/Grant No.: GM30736; GM; NIGMS NIH HHS United States Publishing Model Print

Document type: Evaluation Studies; Journal Article; Research Support,

N.I.H., Extramural; Research Support, U.S. Gov't, Non-P.H.S.; Research

Support, U.S. Gov't, P.H.S.

Languages: ENGLISH

Main Citation Owner: NLM Other Citation Owner: NLM

Record type: MEDLINE; Completed

The gram-negative, purple nonsulfur, facultative photosynthetic bacterium

Rhodobacter capsulatus is a widely used model organism and has

well-developed molecular genetics. In particular, interposon mutagenesis

using selectable gene cartridges is frequently employed for construction of a variety of chromosomal knockout mutants. However,

as the gene cartridges are often derived from antibiotic resistance-conferring genes, their numbers are limited, which

restricts the construction of multiple knockout mutants. In this report.

sacB-5-fluoroorotic acid (5FOA)--pyrE-based bidirectional
selection that

facilitates construction of unmarked chromosomal knockout mutations is

described. The R. capsulatus pyrE gene encoding orotate phosphoribosyl transferase, a key enzyme of the de novo pyrimidine nucleotide biosynthesis pathway, was used as an interposon in a genetic background that is auxotrophic for uracil (Ura-) and hence resistant to 5FOA (5FOA(r)). Although Ura+ selection readily vielded chromosomal allele replacements via homologous recombination, selection for 5FOA(r) to replace pyrE with unmarked alleles was inefficient. To improve the latter step, 5FOA(r) selection was combined with sucrose tolerance selection using a suicide plasmid carrying the Bacillus subtilis sacB gene encoding levansucrase that induces lethality upon exposure to 5% (wt/vol) sucrose in the growth medium. Sucrose-tolerant, 5FOA(r) colonies that were obtained carried chromosomal unmarked mutant alleles of the target gene via double crossovers between the resident pyrE-marked and incoming unmarked alleles. The effectiveness of this double selection was proven by seeking insertion and deletion alleles of helC involved in R. capsulatus cvtochrome c biogenesis, which illustrated the usefulness of this system as a genetic means for facile construction of R. capsulatus unmarked chromosomal mutants. Record Date Created: 20050603 Record Date Completed: 20050811 4/7/14 (Item 3 from file: 154) DIALOG(R) File 154: MEDLINE(R) (c) format only 2009 Dialog. All rts. reserv.

11311703 PMID: 7997171

Five Listeria monocytogenes genes preferentially expressed in infected mammalian cells: plcA, purH, purD, pyrE and an arginine ABC

transporter gene, arpJ.

Klarsfeld A D; Goossens P L; Cossart P

Unite des Interactions Bacteries-Cellules, CNRS URA 1300, Institut

Pasteur, Paris, France.

Molecular microbiology (ENGLAND) Aug 1994, 13 (4) p585-97, ISSN

0950-382X--Print Journal Code: 8712028 Publishing Model Print Document type: Comparative Study; Journal Article; Research Support, Non-U.S. Gov't Languages: ENGLISH Main Citation Owner: NLM Record type: MEDLINE; Completed Listeria monocytogenes is a bacterial pathogen that multiplies within the cytosol of eukaryotic cells. To identify Listeria genes with preferentially intracellular expression (pic genes), a library of Tn917-lac insertion mutants was screened for transcriptional fusions to lacZ with higher expression inside a macrophage-like cell line than in a rich broth Five pic genes with up to 100-fold induction inside cells were identified. Three of them (purH, purD and pyrE) were involved in nucleotide biosynthesis. One was part of an operon encoding an ABC (ATP-binding cassette) transporter for arginine. The corresponding mutants were not affected in intracellular growth, cell-to-cell spread or virulence, for the transporter mutant, whose LD50 after intravenous infection of mice was twofold higher than the wild-type. The fifth gene was plcA, a identified virulence gene that previously encodes phosphatidylinositol-phospholipase C, and is cotranscribed with prfA. a gene encoding a pleiotropic transcriptional activator of known virulence genes. Although plcA expression is known to depend on PrfA, a prfA promoter-lacZ fusion was highly expressed both inside and outside cells. Furthermore, in the presence of cellobiose, a disaccharide recently shown to repress plcA and hly expression, plcA and hly mRNA levels were dramatically reduced without any decrease in the monocistronic prfA mRNA levels. These results demonstrate that virulence gene activation does not depend only on prfA transcript accumulation.

Record Date Created: 19950119

DIALOG(R) File 155: MEDLINE(R) (c) format only 2009 Dialog, All rts, reserv. 07614262 PMID: 6150076 Pyrimidine de novo synthesis during the life cycle of the intraerythrocytic stage of Plasmodium falciparum. Gero A M; Brown G V; O'Sullivan W J Journal of parasitology (UNITED STATES) Aug 1984, 70 (4) p536-41. ISSN 0022-3395--Print Journal Code: 7803124 Publishing Model Print Document type: Journal Article; Research Support, Non-U.S. Gov't Languages: ENGLISH Main Citation Owner: NLM Record type: MEDLINE; Completed The 6 enzymes involved in de novo synthesis of pyrimidines were measured in Plasmodium falciparum isolated by saponin lysis from RBC's nonsynchronized and synchronized in vitro cultures. The total activities were found to be dependent on the stage of the P. falciparum cycle. In parasites isolated from synchronized cultures, the highest activities for all enzymes were found at about 27 hr after synchronization in the late trophozoite stage, or just before schizont formation. Merozoites and ring forms contained little de novo activity. The first enzyme of the pathway, carbamyl phosphate synthetase (CPS-II) preferentially utilized glutamine. Ammonia was a poor substrate, CPS-II was unstable in the absence of the cryoprotectants, dimethylsulfoxide and glycerol. The apparent Km for MgATP--was 3.8 +/- 0.7 mM and the enzyme in all morphological forms of P. falciparum (ring, mature trophozoites and schizonts) was inhibited by UTP. The activity of the fourth enzyme of the pathway, dihydroorotate dehydrogenase, appeared to be linked to the cell's respiratory chain: inhibitors of mammalian electron transport such as cyanide, amvtal, antimycin A, thenoyltrifluoroacetone and ubiquinone analogs also inhibited

the P. falciparum enzyme. The demonstration of the variation of

activity of

pyrimidine enzymes correlates with the increased synthesis of nucleic acids in the late trophozoite stage. These observations provide a basis for the testing of the effectiveness of pyrimidine analogs as potential antimetabolites against various forms of the parasite. Record Date Created: 19850109 Record Date Completed: 19850109 4/7/16 (Item 2 from file: 155) DIALOG(R) File 155: MEDLINE(R) (c) format only 2009 Dialog. All rts. reserv. 05709916 PMTD: 280143 Antiviral action and selectivity of 6-azauridine. Rada B: Dragun M Annals of the New York Academy of Sciences (UNITED STATES) Mar 4 1977. 284 p410-7, ISSN 0077-8923--Print Journal Code: 7506858 Publishing Model Print Document type: Journal Article Languages: ENGLISH Main Citation Owner: NLM Record type: MEDLINE; Completed 6-Azauridine (AzUrd) is a broad-spectrum antimetabolite that inhihits both DNA and RNA virus multiplication. Prior work indicated that several AzUrd-sensitive viruses induced an increase in the level of uridine kinase, and this might explain the selective activity of AzUrd on such viruses. Present studies compared AzUrd sensitive and resistant viruses with respect to their orotic acid pathways by labeling cells with [14C]-orotic acid during the latent period of viral infection. No differences were detected by this method with either vaccinia, Newcastle disease, or vesicular stomatitis viruses. AzUrd inhibits transport of orotic acid into the cell by 30%, while incorporation of orotic acid

into cellular RNA is inhibited by 50% (taking into consideration the 30% already noted) when the highest concentration of antimetabolite is used.

This suggests that, in addition to blocking orotidylic acid decarboxylase,

AzUrd may act on some other site (sites) of action in the inhibition of

virus multiplication.

Record Date Created: 19781227

```
4/7/17
            (Item 3 from file: 155)
DIALOG(R)File 155:MEDLINE(R)
(c) format only 2009 Dialog. All rts. reserv.
05100256
          PMID: 171022
                        Record Identifier: PMC1859231
 Anabolic and androgenic effects of methandrostenolone ("Nerobol")
systematic physical activity in rats.
 Rogozkin V
 British journal of sports medicine (ENGLAND) Jul 1975, 9 (2)
p65-9.
ISSN 0306-3674--Print Journal Code: 0432520
 Publishing Model Print
 Document type: Journal Article
 Languages: ENGLISH
 Main Citation Owner: NLM
 Other Citation Owner: NLM
 Record type: MEDLINE; Completed
 Record Date Created: 19760129
 Record Date Completed: 19760129
 4/7/18
           (Item 1 from file: 370)
DIALOG(R) File 370: Science
(c) 1999 AAAS. All rts. reserv.
00501372
           (USE 9 FOR FULLTEXT)
A Membrane Network for Nutrient Import in Red Cells Infected with the
 Malaria Parasite
Lauer, Sabine A.; Rathod, Pradipsinh K.; Ghori, Nafisa; Haldar,
Kasturi
S. A. Lauer, N. Ghori, K. Haldar, Department of Microbiology and
  Immunology, Stanford University School of Medicine, Stanford, CA
 94305-5402, USA.; P. K. Rathod, Department of Biology, Institute
for
 Biomolecular Studies, The Catholic University of America,
Washington, DC
  20064, USA.
Science Vol. 276 5315 pp. 1122
Publication Date: 5-16-1997 (970516) Publication Year: 1997
Document Type: Journal ISSN: 0036-8075
Language: English
Section Heading: Reports
Word Count: 2537
Abstract:
           The human malaria parasite Plasmodium falciparum exports
an
```

interconnected network of tubovesicular membranes (the TVM) that

extends

```
from the parasite's vacuolar membrane to the periphery of the red
cell.
Here it is shown that extracellular solutes such as Lucifer yellow
enter
the TVM and are delivered to the parasite. Blocking the assembly of
the
network blocked the delivery of exogenous Lucifer yellow,
nucleosides, and
amino acids to the parasite without inhibiting secretion of plasmodial
proteins. These data suggest that the TVM is a transport network that
allows nutrients efficient access to the parasite and could be used to
deliver antimalarial drugs directly into the parasite.
References and Notes:
    Gronowicz, G., Swift, H., Steck, T. L., J. Cell Sci., 171
1984,
 163 :
     Chasis, J. A., Prenant, M., Leung, A., Mohandas, N.,
Blood, 74
 1989, 1112
     Deitsch, K. W., Wellems, T. E., Mol. Biochem. Parasitol., 76
1996,
     Gero, A. M., Kirk, K., Parasitol. Today, 10 1994, 395 ;
5.
     Haldar, K., ibid 393 ;
6.
     Elmendorf, H. G., Haldar, K., J. Cell Biol., 124 1994, 449 ;
     Elford, B. C., Ferguson, D. J. P., Parasitol. Today, 9 1993,
7.
80 ;
8. Lauer, S., Ghori, N., Haldar, K., Proc. Natl. Acad. Sci.
 92 1995, 9181 PPMP-treated cells were prepared by incubating
purified
 schizonts (36 to 48 hours old) with a 20-fold excess of uninfected
  cells for 12 to 16 hours in RPMI 1640 medium containing 10% human
serum
  and 5 (mu) M PPMP. This treatment produced a new generation of
 PPMP-treated infected red cells that were arrested in the TVM from
the
 onset of ring development. Any residual schizonts were removed by a
 second Percoll gradient. ;
     Haldar, K., de Amorin, A. F., Cross, G. A. M., J. Cell
Biol., 108
 1989, 2183 ;
10.
      Haldar, K., Uyetake, L., Mol. Biochem. Parasitol., 50 1992,
161
11.
     Pouvelle, B., Gormley, J. A., Taraschi, T. F., ibid., 66
1994.
12. To examine the uptake of LY after removal of PPMP, we washed
```

cells in

culture medium, allowed them to mature under normal growth conditions for

24 hours, and subsequently incubated them with LY (B9) . To examine the

effect of PPMP on the LY channel or transporter activity in the infected

 $\ensuremath{\operatorname{red}}$ cell membrane, we incubated trophozoite-and schizont-infected $\ensuremath{\operatorname{red}}$

cells with LY (B9) in the presence or absence of 5 $\,$ (mu) M PPMP. The dye

 $\mbox{\tt could}$ be detected in both treated and control cells, indicating that $\mbox{\tt PPMP}$

does not block the channel or transporter activity at the infected red

cell membrane. ;

13. Gero, A. M., Bugledich, E. M. A., Paterson, A. R. P., Jamieson,

G. P., Mol. Biochem. Parasitol., 27 1988, 159 ;

14. PPMP-treated and control cells were adjusted to 2 x 10.sup(7) cells/ml and incubated with a 1 (mu) Ci/ml concentration of [.sup(3)H]adenosine (34.5 Ci/mmol at a final concentration 29 nM) or [.sup(3)H]thymidine (2 Ci/mmol at a final concentration of 500 nM)

phosphate-buffered saline (PBS). Because of its low specific activity.

the extracellular concentration of [.sup(3)H]thymidine was 16 times

high as that of adenosine. For the accumulation of orotic acid, infected

erythrocytes treated with 5 $\,$ (mu) M PPMP (10 to 20% parasitemia) were

washed free of serum and adjusted to 5 $\,\mathrm{x}\,$ 10.sup(9) cells/ml in RPMT

1640. Transport was initiated by mixing equal volumes (330 (mu) 1) of

the cell suspension and [.sup(3)H]orotic acid at a concentration of 0.26

(mu) Ci/ml (13 Ci/mmol at a final concentration of 10 nM). For the accumulation of glutamate, control and PPMP-treated infected red cells

were adjusted to 2.5 x $10.\sup(8)$ cells/ml and incubated with [. $\sup(3)$ H]glutamate at a concentration of 1 (mu) Ci/ml (53 Ci/mmol at a

final concentration of 19 nM) in PBS. For all accumulation assays, the $\,$

cells were collected by centrifugation through a layer of dibutylphthalate, and the cell pellets lysed, bleached, neutralized, and

counted. PPMP treatment had no effect on the uptake of any radiolabeled

compound into uninfected red cells. The incorporation of adenosine and $% \left(1\right) =\left(1\right) +\left(1\right) +\left($

```
orotic acid into nucleic acids was measured as follows. Infected
 red cells treated with PPMP (5 (mu) M) for 12 hours and their
 corresponding controls were seeded into microtiter plates at 2%
 hematocrit and 5% parasitemia. The cells were incubated in RPMI 1640
 containing 10% human serum and 0.86 (mu) Ci/well (11 nM) of
  [.sup(3)H]adenosine or 0.125 (mu) Ci/well (48 nM) of
[.sup(3)H]orotic
 acid in the presence or absence of (5 (mu) M) PPMP for 24 hours at
  37.Deq.C. All incorporation assays were carried out in the absence
 NBMPR. For glutamate, infected red cells treated for 12 hours with
PPMP
  (5 (mu) M) and their corresponding controls were incubated for 24
hours
 at 37.Deg.C in RPMI 1640 lacking glutamate and supplemented with 2%
human
 serum and [.sup(3)H]qlutamic acid (20 (mu) Ci/ml, 377 nM).;
      Crary, J. L., Haldar, K., Mol. Biochem. Parasitol., 53
1992, 185
16.
      Elford, B. C., Cowman, G. M., Ferguson, D. J. P., Biochem.
  308 1995, 361 ;
     Elmendorf, H. G., Bangs, J. D., Haldar, K., Mol. Biochem.
 Parasitol., 52 1992, 215
      Rathod, P. K., Khatri, A., Hubbert, T., Milhous, W. K.,
 Antimicrob. Agents Chemother., 33 1989, 1090 ;

    Determination of the IC.inf(50) of 5-FO in TVM-arrested cells by

 hypoxanthine incorporation was carried out by plating infected red
cells
  in microtiter dishes at 1% hematocrit and 1% parasitemia.
  [.sup(3)H]Hypoxanthine (0.5 (mu) Ci/well) was added, and the cells
were
  incubated for another 24 hours and then harvested on glass fiber
filters.
  (B8) . Because PPMP reduces the accumulation of exogenous
  [.sup(3)H]hypoxanthine (but does not inhibit the parasite's
machinery for
```

determined at 0, 0.03, 0.3, and 3.3 (mu) M PPMP. At 0 to 0.3 (mu) M PPMP, the IC.inf(50) was 6.0~x~10.sup(-8) M. At 3.3 (mu) M PPMP (which

nucleic acid synthesis), a separate IC.inf(50) plot for 5-FO was

corresponds to complete inhibition of the SSS and the TVM), the IC.inf(50) of 5-PO was 7.5 x 10.sup(-7) M. As expected, saponin abrogated the effects on nucleoside uptake. Determination of the IC.inf(50) in TVM-arrested cells by Giemsa staining was carried out with

infected red cells at 2% parasitemia and 5% hematocrit. Cells were subsequently washed free of both PPMP and 5-FO and the parasites were

allowed to grow in RPMI 1640 for another 48 hours, at which time

parasitemia was determined by Giemsa staining. The IC.inf(50) of doxycline (in the absence or presence of PPMP) was 1 x 10.sup(-6) Μ. These experiments were carried out as those described for 5-FO.; 20. Kirk, K., Horner, H. A., Elford, B. C., Ellory, J. C., Newbold, C., J. Biol. Chem., 269 1994, 3339 Pouvelle, B., et.al. Nature, 353 1991, 73 ; Taraschi, T. F., Nicholas, E., Parasitol. Today, 10 1994, 399 ; 23. Desai, S. A., Krogstad, D. J., McClesky, E. W., Nature, 362 1993, 643 ; 24. Swedlow, J. R., Sedat, J. W., Agard, D. A., Cell, 9 1993, 97 ; 25. We thank W. l. Li, J. McBride, D. Taylor, and R. Coppel for antibodies to the 45-kD cleft protein, Expl, HRP1, HRP2, and PfEMP2; A. A. Holder for a knob-forming FCB strain of P. falciparum; R. R. Kopito and S. Mayor for comments on the manuscript; and S. Palmieri and J. VanWye for assistance with the Delta Vision microscope and work station. Supported by NIH grants (AI26670 and AI39071 to K.H., AI26912 and AI01112 to P.K.R.), the MacArthur Foundation (S.L.), and Burroughs Wellcome Awards (New Investigator and New Initiatives in Malaria to K.H.). 4/7/19 (Item 1 from file: 399) DIALOG(R) File 399:CA SEARCH(R) (c) 2009 American Chemical Society. All rts. reserv. 145477770 CA: 145(24)477770v PATENT Facilitated transport of bisphosphonates by vitamin C, and use of bisphosphonate-ascorbic acid conjugates for treatment of joint and bone diseases INVENTOR(AUTHOR): Kraus, Virginia Byers; McNulty, Amy Lynn; Toone, Eric John LOCATION: USA ASSIGNEE: Duke University PATENT: PCT International ; WO 2006116057 A2 DATE: 20061102 APPLICATION: WO 2006US15051 (20060421) *US 2005PV673527 (20050421) PAGES: 65pp. CODEN: PIXXD2 LANGUAGE: English PATENT CLASSIFICATIONS: IPCR/8 + Level Value Position Status Version Action Source Office: A I F B 20060101 C07F-0009/58 DESIGNATED COUNTRIES: AE; AG; AL; AM; AT; AU; AZ; BA; BB; BG; BR; BZ: CA: CH: CN: CO: CR: CU: CZ: DE: DK: DM: DZ: EC: EE: EG: ES: FI:

GB; GD;

```
GE; GH; GM; HR; HU; ID; IL; IN; IS; JP; KE; KG; KM; KN; KP; KR; KZ;
LC: LK:
LR; LS; LT; LU; LV; MA; MD; MG; MK; MN; MW; MX; MZ; NA; NG; NI;
NO: NZ:
OM; PG; PH; PL; PT; RO; RU; SC; SD; SE; SG; SK; SL; SM; SY; TJ; TM;
TN; TR;
TT; TZ; UA; UG; US; UZ; VC; VN; YU; ZA DESIGNATED REGIONAL: AT; BE;
BG; CH
; CY; CZ; DE; DK; EE; ES; FI; FR; GB; GR; HU; IE; IS; IT; LT; LU; LV;
NL; PL; PT; RO; SE; SI; SK; TR; BF; BJ; CF; CG; CI; CM; GA; GN; GQ;
GW: ML:
MR; NE; SN; TD; TG; BW; GH; GM; KE; LS; MW; MZ; NA; SD; SL; SZ; TZ;
UG; ZM;
ZW; AM; AZ; BY; KG; KZ; MD; RU; TJ; TM
 SECTION:
    CA263005 Pharmaceuticals
    CA201XXX Pharmacology
    CA213XXX Mammalian Biochemistry
  IDENTIFIERS: bisphosphonate ascorbic acid conjugate cartilage uptake
    gastrointestinal absorption, joint bone disease treatment
    bisphosphonate ascorbic acid conjugate
  DESCRIPTORS:
Bone, disease...
    abnormally increased bone turnover; facilitated transport of
    bisphosphonates by vitamin C, and use of bisphosphonate-ascorbic
acid
    conjugates for treatment of joint and bone diseases
Disease, animal...
    arthropathy; facilitated transport of bisphosphonates by vitamin
    use of bisphosphonate-ascorbic acid conjugates for treatment of
ioint
    and bone diseases
Peptides, biological studies... Glycosaminoglycans, biological
Imaging agents... Nucleic acids... Enzyme inhibitors... Antitumor
agents...
    conjugates with ascorbic acids or analogs; facilitated transport
\circ f
    bisphosphonates by vitamin C, and use of bisphosphonate-ascorbic
acid
    conjugates for treatment of joint and bone diseases
Cartilage, disease... Tendon... Synovial membrane, disease...
    degeneration; facilitated transport of bisphosphonates by vitamin
С,
    and use of bisphosphonate-ascorbic acid conjugates for treatment
of
    joint and bone diseases
Epithelium ...
    digestive tract, transport across; facilitated transport of
    bisphosphonates by vitamin C, and use of bisphosphonate-ascorbic
```

acid

```
conjugates for treatment of joint and bone diseases
Joint, anatomical ...
    disease, degeneration; facilitated transport of bisphosphonates by
    vitamin C, and use of bisphosphonate-ascorbic acid conjugates for
    treatment of joint and bone diseases
Joint, anatomical...
    disease; facilitated transport of bisphosphonates by vitamin C,
    of bisphosphonate-ascorbic acid conjugates for treatment of joint
and
    bone diseases
Biological transport ...
    drug; facilitated transport of bisphosphonates by vitamin C, and
use of
    bisphosphonate-ascorbic acid conjugates for treatment of joint
and bone
    diseases
Calcification...
    ectopic; facilitated transport of bisphosphonates by vitamin C,
and use
    of bisphosphonate-ascorbic acid conjugates for treatment of joint
and
    bone diseases
Joint, anatomical... Cartilage... Synovial membrane...
    enhancing joint tissue synthesis; facilitated transport of
    bisphosphonates by vitamin C, and use of bisphosphonate-ascorbic
acid
    conjugates for treatment of joint and bone diseases
Digestive tract...
    epithelium, transport across; facilitated transport of
bisphosphonates
    by vitamin C, and use of bisphosphonate-ascorbic acid conjugates
    treatment of joint and bone diseases
Human... Diphosphonates... Osteoporosis... Antiosteoporotic agents...
... Osteoarthritis... Rheumatoid arthritis... Antiarthritics...
Periodontium, disease... Multiple myeloma...
    facilitated transport of bisphosphonates by vitamin C, and use of
    bisphosphonate-ascorbic acid conjugates for treatment of joint
and bone
   diseases
Bone, disease . . .
    fracture; facilitated transport of bisphosphonates by vitamin C,
and
    use of bisphosphonate-ascorbic acid conjugates for treatment of
ioint
    and bone diseases
Transport proteins ...
    GLUT-1 (glucose transporter 1), GLUTs-mediated DHA transport in
human
    chondrocytes is regulated by hypoxia; facilitated transport of
```

bisphosphonates by vitamin \mathbf{C}_{\bullet} and use of bisphosphonate-ascorbic ac

Transport proteins...

 $\ensuremath{\mathsf{GLUT-3}}$ (glucose transporter 3), GLUTs-mediated DHA transport in human

chondrocytes is regulated by hypoxia; facilitated transport of bisphosphonates by vitamin C, and use of bisphosphonate-ascorbic

Hypoxia, animal...

GLUTs-mediated DHA transport in human chondrocytes is regulated by hypoxia; facilitated transport of bisphosphonates by vitamin C, and use

of bisphosphonate-ascorbic acid conjugates for treatment of j

Neoplasm...

humoral hypercalcemia of malignancy; facilitated transport of bisphosphonates by vitamin C, and use of bisphosphonate-ascorbic acid

conjugates for treatment of joint and bone diseases

Drug delivery systems...

injections, intraarticular; facilitated transport of bisphosphonates by

vitamin C, and use of bisphosphonate-ascorbic acid conjugates for treatment of joint and bone diseases

Disease, animal...

joint degeneration; facilitated transport of bisphosphonates by vitamin

C, and use of bisphosphonate-ascorbic acid conjugates for treatment of

joint and bone diseases

Bone, neoplasm...

metastasis; facilitated transport of bisphosphonates by vitamin \mathbf{C} , and

use of bisphosphonate-ascorbic acid conjugates for treatment of $\ensuremath{\operatorname{\textsc{ioint}}}$

and bone diseases

Stereochemistry...

of ascorbic acid by chondrocytes; facilitated transport of bisphosphonates by vitamin C, and use of bisphosphonate-ascorbic acid

conjugates for treatment of joint and bone diseases

Bone, disease...

Paget's; facilitated transport of bisphosphonates by vitamin C, and use

of bisphosphonate-ascorbic acid conjugates for treatment of joint and $% \left(1\right) =\left(1\right) +\left(1$

bone diseases

Transport proteins ...

SVCT2, chondrocyte transport of vitamin C mediated by; facilitated transport of bisphosphonates by vitamin C, and use of bisphosphonate-ascorbic acid conjugates for treatment of joint

and bone

disease

SVCT2-mediated transport of vitamin C into chondrocytes: facilitated transport of bisphosphonates by vitamin C, and use of bisphosphonate-ascorbic acid conjugates for treatment of joint and bone disea Tooth, disease ... tooth loss; facilitated transport of bisphosphonates by vitamin use of bisphosphonate-ascorbic acid conjugates for treatment of ioint and bone diseases Biological transport ... uptake, carrier-mediated, SVCT2-mediated transport of vitamin C into chondrocytes; facilitated transport of bisphosphonates by vitamin C. and use of bisphosphonate-ascorbic acid conjugates for treatme CAS REGISTRY NUMBERS: 50-81-7 biological studies, facilitated transport of bisphosphonates by vitamin C, and use of bisphosphonate-ascorbic acid conjugates for treatment of joint and bone diseases 7440-23-5 biological studies, vitamin C transport dependence on; facilitated transport of bisphosphonates by vitamin C, and use of bisphosphonate-ascorbic acid conjugates for treatment of joint and bone diseases 50-81-7D conjugates with bisphosphonates, facilitated transport of bisphosphonates by vitamin C, and use of bisphosphonate-ascorbic acid conjugates for treatment of joint and bone diseases 913824-02-9 facilitated transport of bisphosphonates by vitamin C, and use of bisphosphonate-ascorbic acid conjugates for treatment of joint and bone diseases 490-83-5 GLUTs-mediated DHA transport in human chondrocytes is regulated by hypoxia; facilitated transport of bisphosphonates by vitamin C, and

use of bisphosphonate-ascorbic acid conjugates for treatment of

Chondrocyte...

ioint.

4/7/20

and bone diseases

DIALOG(R)File 399:CA SEARCH(R)
(c) 2009 American Chemical Society. All rts. reserv.

144206477 CA: 144(12)206477m JOURNAL

(Item 2 from file: 399)

Protein and cDNA sequences of a novel Lactococcus lactis orotate transporter and use as selection markers LOCATION: Den.

JOURNAL: IP.com J. (IP.com Journal) DATE: 2005 VOLUME: 5 NUMBER:

PAGES: 23 CODEN: TJPOBX JSSN: 1533-0001 JSSN: JPCOM000124929D LANGUAGE: English PUBLISHER: IP.com, Inc. SECTION:

CA203001 Biochemical Genetics CA206XXX General Biochemistry

CA209XXX Biochemical Methods

CA210XXX MICROBIAL, ALGAL, AND FUNGAL BIOCHEMISTRY

IDENTIFIERS: sequence Lactococcus orotate transport protein gene selection marker DESCRIPTORS:

Escherichia coli... Bacillus subtilis...

expression host; protein and cDNA sequences of novel Lactococcus

orotate transporter and use as selection markers

Transport proteins ...

orotate transporter; protein and cDNA sequences of novel

Lactococcus

lactis orotate transporter and use as selection markers Lactococcus lactis... Protein sequences... cDNA sequences... Molecular

cloning ... Selection markers ... protein and cDNA sequences of novel Lactococcus lactis orotate

transporter and use as selection markers Gene, animal ...

ysbC, for orotate transporter; protein and cDNA sequences of novel Lactococcus lactis orotate transporter and use as selection markers

CAS REGISTRY NUMBERS:

875602-72-5P amino acid sequence; protein and cDNA sequences of novel Lactococcus lactis orotate transporter and use as selection markers

65-86-1D analog, protein and cDNA sequences of novel Lactococcus lactis

orotate transporter and use as selection markers

875602-71-4P nucleotide sequence; protein and cDNA sequences of a novel

Lactococcus lactis orotate transporter and use as selection markers

703-95-7 protein and cDNA sequences of novel Lactococcus lactis orotate

transporter and use as selection markers

(Item 3 from file: 399) 4/7/21 DIALOG(R) File 399:CA SEARCH(R)

(c) 2009 American Chemical Society. All rts. reserv.

143242974 CA: 143(14)242974; PATENT

```
Protein and cDNA sequences of a novel Lactococcus lactis orotate
  transporter and use as selection markers
  INVENTOR(AUTHOR): Martinussen, Jan; Defoor, Els Marie Celine
  LOCATION: Den.
  ASSIGNEE: Novozymes A/S
  PATENT: PCT International; WO 200578106 Al DATE: 20050825
  APPLICATION: WO 2005DK92 (20050211) *DK 2004227 (20040213)
  PAGES: 58 pp. CODEN: PIXXD2 LANGUAGE: English
  PATENT CLASSIFICATIONS:
    CLASS: C12N-015/65A; C12N-001/21B; C12N-015/31B; C12N-015/64B;
C12N-015/70B; C12N-015/74B; C12N-015/75B
  DESIGNATED COUNTRIES: AE; AG; AL; AM; AT; AU; AZ; BA; BB; BG; BR;
BW; BY;
BZ; CA; CH; CN; CO; CR; CU; CZ; DE; DK; DM; DZ; EC; EE; EG; ES; FI;
GB; GD;
GE; GH; GM; HR; HU; ID; IL; IN; IS; JP; KE; KG; KP; KR; KZ; LC; LK;
LR: LS:
LT: LU: LV: MA: MD: MG: MK: MN: MW: MX: MZ: NA: NI: NO: NZ: OM: PG:
PH; PL;
PT; RO; RU; SC; SD; SE; SG; SK; SL; SY; TJ; TM; TN; TR; TT; TZ; UA;
UG; US;
UZ; VC; VN; YU; ZA; ZM; ZW DESIGNATED REGIONAL: BW; GH; GM; KE; LS;
MW; MZ
; NA; SD; SL; SZ; TZ; UG; ZM; ZW; AM; AZ; BY; KG; KZ; MD; RU; TJ; TM;
AT:
BE; BG; CH; CY; CZ; DE; DK; EE; ES; FI; FR; GB; GR; HU; IE; IS; IT;
LT: LU:
MC; NL; PL; PT; RO; SE; SI; SK; TR; BF; BJ; CF; CG; CI; CM; GA; GN;
GO: GW:
ML; MR; NE; SN; TD; TG
  SECTION:
    CA203001 Biochemical Genetics
    CA206XXX General Biochemistry
    CA209XXX Biochemical Methods
    CA210XXX MICROBIAL, ALGAL, AND FUNGAL BIOCHEMISTRY
  IDENTIFIERS: sequence Lactococcus orotate transport protein gene
    selection marker
  DESCRIPTORS:
Microorganism... Eubacteria... Firmicutes... Lactobacillus...
Bacillus(bacterium genus) ... Escherichia ...
    as expression host; protein and cDNA sequences of novel
Lactococcus
    lactis orotate transporter and use as selection markers
    for orotate transporter; protein and cDNA sequences of novel
    Lactococcus lactis orotate transporter and use as selection
markers
Transport proteins ...
```

orotate; protein and cDNA sequences of novel Lactococcus lactis transporter and use as selection markers Lactococcus lactis... Protein sequences... cDNA sequences... Molecular

orotate

```
cloning... Selection markers... Promoter(genetic element)... Genetic
vectors... Plasmid vectors... Mutation...
```

protein and cDNA sequences of novel Lactococcus lactis orotate transporter and use as selection markers $\,$

Gene, microbial...
pvrD; protein and cDNA sequences of novel Lactococcus lactis

orotate

transporter and use as selection markers

Gene, microbial...

 $\ensuremath{\operatorname{pyrDa;}}$ protein and cDNA sequences of novel Lactococcus lactis orotate

transporter and use as selection markers

Gene, microbial...

 $\ensuremath{\operatorname{pyrDb}};$ protein and cDNA sequences of novel Lactococcus lactis orotate

transporter and use as selection markers Gene, microbial...

 $\ensuremath{\operatorname{\textsc{pyrK}}};$ protein and cDNA sequences of novel Lactococcus lactis orotate

transporter and use as selection markers

CAS REGISTRY NUMBERS:

863466-31-3P amino acid sequence; protein and cDNA sequences of novel Lactococcus lactis orotate transporter and use as selection markers

65--86--1D analog, protein and cDNA sequences of novel Lactococcus lactis

orotate transporter and use as selection markers

66-22-8 biological studies, protein and cDNA sequences of novel Lactococcus lactis orotate transporter and use as selection markers

863466-30-2P nucleotide sequence; ;protein and cDNA sequences of novel

Lactococcus lactis orotate transporter and use as selection markers $% \left(1\right) =\left(1\right) +\left(1\right)$

703-95-7 289-95-2 155-54-4 protein and cDNA sequences of novel Lactococcus lactis orotate transporter and use as selection markers

863467-19-0 863467-20-3 863467-21-4 863467-22-5 863467-23-6 863467-24-7 863467-25-8 863467-26-9 863467-27-0 863467-28-1 863467-29-2 863467-30-5 863467-31-6 863467-32-7 863467-38-3 863467-34-9 863467-35-0 863467-36-1 863467-37-2 863467-38-3 unclaimed nucleotide sequence; protein and cDNA sequences of a

Lactococcus lactis orotate transporter and use as selection markers

4/7/22 (Item 4 from file: 399)
DIALOG(R)File 399:CA SEARCH(R)

(c) 2009 American Chemical Society. All rts. reserv.

141406805 CA: 141(25)406805s PATENT

```
Cloning of orotate-phosphoribosyl transferase gene Ura5 from P.
pastoris
  and use thereof as a new selection marker for stable genetic
integration
 in veast
 INVENTOR (AUTHOR): Nett, Juergen H.
 LOCATION: USA
 PATENT: U.S. Pat. Appl. Publ.; US 20040229306 Al DATE: 20041118
 APPLICATION: US 454125 (20030603) *US PV471435 (20030516)
 PAGES: 38 pp. CODEN: USXXCO LANGUAGE: English
 PATENT CLASSIFICATIONS:
    CLASS: 435069100; C12N-009/10A; C12N-001/18B; C07H-021/04B;
C12N-015/74B
 SECTION:
    CA203003 Biochemical Genetics
    CA207XXX Enzymes
    CA210XXX MICROBIAL, ALGAL, AND FUNGAL BIOCHEMISTRY
  IDENTIFIERS: sequence orotate phosphoribosyltransferase gene URA5
Pichia
    transformation selection marker, Pichia SEC65 SCS7 gene fragment
    sequence
 DESCRIPTORS:
Protein sequences... DNA sequences... Pichia pastoris...
Transformation, genetic... Cytokines... Blood-coagulation factors...
Insulin-like growth factor-binding proteins... \alpha-Fetoproteins...
Antibodies and Immunoglobulins... Molecular cloning...
    cloning of orotate-phosphoribosyl transferase gene Ura5 from P.
    pastoris and use thereof as a new selection marker for stable
genetic
    integration in yeast
Repetitive DNA ...
    direct, flanking Ura3 or Ura5 gene; cloning of
orotate-phosphoribosvl
    transferase gene Ura5 from P. pastoris and use thereof as a new
    selection marker for stable genetic integration in yeast
Glycosylation...
    enzyme, disruption of; cloning of orotate-phosphoribosyl
transferase
    gene Ura5 from P. pastoris and use thereof as a new selection
marker
    for stable genetic integration in yeast
Antibodies and Immunoglobulins...
    fragments, antigen-binding; cloning of orotate-phosphoribosyl
    transferase gene Ura5 from P. pastoris and use thereof as a new
    selection marker for stable genetic integration in yeast
```

from P. pastoris and use thereof as a new selection marker for stable genetic integration in yeast

gene knock-out; cloning of orotate-phosphoribosyl transferase

Proteins...

gene Ura5

Gene targeting ...

gene SCS7; cloning of orotate-phosphoribosyl transferase gene Ura5 from

P. pastoris and use thereof as a new selection marker for stable genetic integration in yeast

Proteins...

gene SEC65; cloning of orotate-phosphoribosyl transferase gene $\mbox{Ura5}$

from P. pastoris and use thereof as a new selection marker for stable $% \left(1\right) =\left(1\right) +\left(1$

genetic integration in yeast

Immunoglobulin receptors...

IgE, sol., α -chain; cloning of orotate-phosphoribosyl transferase gene Ura5 from P. pastoris and use thereof as a new selection marker

for stable genetic integration in yeast

Antibodies and Immunoglobulins ...

IgG, fragment; cloning of orotate-phosphoribosyl transferase gene IIra5

from P. pastoris and use thereof as a new selection marker for stable $% \left(1\right) =\left(1\right) +\left(1$

genetic integration in yeast

Antibodies and Immunoglobulins...

IgG; cloning of orotate-phosphoribosyl transferase gene Ura5 from
P.

pastoris and use thereof as a new selection marker for stable genetic

integration in veast

Antibodies and Immunoglobulins ...

IgM; cloning of orotate-phosphoribosyl transferase gene $\operatorname{Ura5}$ from $\operatorname{P.}$

pastoris and use thereof as a new selection marker for stable $\ensuremath{\mathsf{genetic}}$

integration in yeast

Recombination, genetic ...

integration; cloning of orotate-phosphoribosyl transferase gene Ura5

from P. pastoris and use thereof as a new selection marker for stable $% \left(1\right) =\left(1\right) +\left(1$

genetic integration in yeast

Chemokines...

macrophage inflammatory protein 3; cloning of

orotate-phosphoribosyl

transferase gene Ura5 from P. pastoris and use thereof as a new selection marker for stable genetic integration in yeast

Transport proteins...

nucleotide sugar-transporting, gene disruption; cloning of orotate-phosphoribosyl transferase gene Ura5 from P. pastoris and

thereof as a new colection marker for stable con-

thereof as a new selection marker for stable genetic integration in yea $\,$

Gene, microbial ...

OCH1, knockout of; cloning of orotate-phosphoribosyl transferase gene

```
Ura5 from P. pastoris and use thereof as a new selection marker
for
    stable genetic integration in yeast
Fusion proteins(chimeric proteins)...
    of orotate phosphoribosyltransferase and SEC65p and SCS7p;
cloning of
    orotate-phosphoribosyl transferase gene Ura5 from P. pastoris and
use
    thereof as a new selection marker for stable genetic integra
Enzymes, processes...
    Phosphomannosidase, gene disruption; cloning of
orotate-phosphoribosyl
    transferase gene Ura5 from P. pastoris and use thereof as a new
    selection marker for stable genetic integration in yeast
Plasmid vectors...
    pJN266, disruption vector contq. Ura3; cloning of
    orotate-phosphoribosyl transferase gene Ura5 from P. pastoris and
    thereof as a new selection marker for stable genetic integration
in
    veast
Plasmid vectors...
    pJN395, disruption vector contq, Ura5; cloning of
    orotate-phosphoribosyl transferase gene Ura5 from P. pastoris and
use
    thereof as a new selection marker for stable genetic integration
in
    veast
Plasmid vectors...
    pJN396, disruption vector contq. Ura5; cloning of
    orotate-phosphoribosyl transferase gene Ura5 from P. pastoris and
1186
    thereof as a new selection marker for stable genetic integration
in
    veast
Plasmid vectors...
    pJN398, disruption vector contg. Ura5; cloning of
    orotate-phosphoribosyl transferase gene Ura5 from P. pastoris and
use
    thereof as a new selection marker for stable genetic integration
in
    yeast
Plasmid vectors...
    pJN407, disruption vector contq. Ura5 and UDP-GlcNAc transporter
gene;
    cloning of orotate-phosphoribosyl transferase gene Ura5 from P.
    pastoris and use thereof as a new selection marker for stable gen
Pichia pastoris... Pichia finlandica... Pichia trehalophila... Pichia
```

kodamae... Pichia membranaefaciens... Pichia opuntiae... Pichia thermotolerans... Pichia salictaria... Pichia quercuum... Pichia

Yamadazyma stipite... Pichia methanolica... Pichia... Saccharomyces

pijperi...

```
cerevisiae... Kluyveromyces lactis... Pichia angusta...
Kluvveromvces...
Candida albicans... Aspergillus nidulans... Aspergillus niger...
Aspergillus oryzae... Trichoderma reesei... Chrysosporium
lucknowense...
Fusarium ... Fusarium graminearum ... Fusarium venenatum ... Neurospora
crassa
. . .
    recombinant host; cloning of orotate-phosphoribosyl transferase
gene
    Ura5 from P. pastoris and use thereof as a new selection marker
for
    stable genetic integration in veast
Gene, microbial ...
    SCS7; cloning of orotate-phosphoribosyl transferase gene Ura5
from P.
    pastoris and use thereof as a new selection marker for stable
genetic
    integration in yeast
Gene, microbial ...
    SEC65; cloning of orotate-phosphoribosyl transferase gene Ura5
from P.
    pastoris and use thereof as a new selection marker for stable
genetic
    integration in veast
Proteins...
    therapeutic; cloning of orotate-phosphoribosyl transferase gene
Ura5
    from P. pastoris and use thereof as a new selection marker for
stable
    genetic integration in yeast
Transport proteins ...
    UDP-N-acetylglucosamine transporting, gene for; cloning of
    orotate-phosphoribosyl transferase gene Ura5 from P. pastoris and
11SP
    thereof as a new selection marker for stable genetic integration
in vea
Transport proteins ...
    UDP-N-acetylglucosamine-transporting, gene disruption; cloning of
    orotate-phosphoribosyl transferase gene Ura5 from P. pastoris and
use
    thereof as a new selection marker for stable genetic integration
```

Gene,microbial...
 URA5; cloning of orotate-phosphoribosyl transferase gene Ura5
from P.
 pastoris and use thereof as a new selection marker for stable
genetic

URA3; cloning of orotate-phosphoribosyl transferase gene Ura5

pastoris and use thereof as a new selection marker for stable

Gene, microbial ...

integration in veast

from P.

genetic

integration in yeast Annexins... V, fusion product; cloning of orotate-phosphoribosyl transferase gene Ura5 from P. pastoris and use thereof as a new selection marker for stable genetic integration in yeast CAS REGISTRY NUMBERS: 791704-69-3 791704-71-7 791704-73-9 amino acid sequence; cloning of orotate-phosphoribosyl transferase gene Ura5 from P. pastoris and use thereof as a new selection marker for stable genetic integration in yeast 66-22-8 biological studies, auxotrophy for; cloning of orotate-phosphoribosyl transferase gene Ura5 from P. pastoris and use thereof as a new selection marker for stable genetic integration

yeast 11096-26-7P 9039-53-6P 97501-92-3P 9035-81-8P 62229-50-9P

9034-39-3P
86090-08-6P 244019-42-9P 205944-50-9P 9041-92-3P 9025-64-3P
cloning of orotate-phosphoribosyl transferase gene Ura5 from P.
pastoris and use thereof as a new selection marker for stable

genetic integration in veast

in

9032-92-2 37211-66-8 9013-05-2 9055-06-5 9054-49-3 9031-68-9 321976-25-4 gene disruption; cloning of orotate-phosphoribosyl transferase gene Ura5 from P. pastoris and use thereof as a new selection marker for stable genetic integration in yeast 9030-25-5 gene Ura5; cloning of orotate-phosphoribosyl transferase gene

Ura5 from P. pastoris and use thereof as a new selection marker for

stable genetic integration in yeast

9001-91-6P kringle domain, of human; cloning of orotate-phosphoribosyl

transferase gene Ura5 from P. pastoris and use thereof as a new selection marker for stable genetic integration in yeast 791704-67-1 791704-68-2 791704-70-6 791704-72-8 nucleotide

/91/04-6/-1 /91/04-68-2 /91/04-70-6 /91/04-72-8 nucleotide sequence; cloning of orotate-phosphoribosyl transferase gene Ura5 from P.

cloning of orotate-phosphoribosyl transferase gene Ura5 from P pastoris and use thereof as a new selection marker for stable genetic

integration in yeast

703-95-7 resistance to; cloning of orotate-phosphoribosyl transferase gene

Ura5 from P. pastoris and use thereof as a new selection marker for

stable genetic integration in yeast 791705-65-2 791705-66-3 791705-67-4 791705-68-5 791705-69-6

```
791705-70-9 791705-71-0 791705-72-1 791705-73-2 791705-74-3
   791705-75-4 791705-76-5 791705-77-6 791705-78-7 791705-79-8
   791705-80-1 791705-81-2 791705-82-3 791705-83-4 791705-84-5
   791705-85-6 791705-86-7 791705-87-8 791705-88-9 791705-89-0
   unclaimed nucleotide sequence; cloning of orotate-phosphoribosyl
   transferase gene Ura5 from P. pastoris and use thereof as a new
    selection marker for stable genetic integration in yeast
791705-50-5 791705-51-6 791705-52-7 791705-53-8 791705-54-9
   791705-55-0 791705-56-1 791705-57-2 791705-58-3 791705-59-4
   791705-60-7 791705-61-8 791705-62-9 791705-63-0 791705-64-1
   unclaimed protein sequence; cloning of orotate-phosphoribosyl
   transferase gene Ura5 from P. pastoris and use thereof as a new
    selection marker for stable genetic integration in yeast
 4/7/23
           (Item 5 from file: 399)
DIALOG(R)File 399:CA SEARCH(R)
(c) 2009 American Chemical Society. All rts. reserv.
              CA: 100(17)134444g TECHNICAL REPORT
 Effect of some derivatives of pyrimidine and purine on RNA transfer
from
 hepatocyte nuclei in a cell-free system
 AUTHOR(S): Zhemkova, L. N.; Porollo, V. I.; Nesterova, S. M.;
Komar, V.
E.; Khanson, K. P.
  LOCATION: Tsentr. Nauchno-Issled. Rentgeno-Radiol. Inst.,
Leningrad, USSR
 JOURNAL: Deposited Doc. DATE: 1982 NUMBER: VINITI 541-83 PAGES:
16 pp.
 CODEN: D8DEP2 LANGUAGE: Russian AVAIL: VINITI
 SECTION:
   CA106001 General Biochemistry
  IDENTIFIERS: RNA transport liver regeneration stimulator, kinetin
liver
    regeneration RNA transport, orotate liver regeneration RNA
transport.
   methyluracil liver regeneration RNA transport
  DESCRIPTORS:
Regeneration, biological...
    of liver, RNA efflux from hepatocyte nuclei in, kinetin and
potassium
   orotate effect on
Biological transport, efflux...
    of RNA, from hepatocyte nuclei, kinetin and potassium orotate
effect on
Cell nucleus...
   RNA transfer from, kinetin and potassium orotate stimulation of,
    regeneration in relation to
Liver, hepatocyte, metabolism...
   RNA transfer from nuclei of, kinetin and potassium orotate
stimulation
```

of, regeneration in relation to Ribonucleic acids ... transfer of, from hepatocyte nuclei, kinetin and potassium orotate effect on CAS REGISTRY NUMBERS: 626-48-2 liver regeneration stimulation by, RNA transport in relation to 525-79-1 24598-73-0 RNA efflux from nuclei enhancement by, liver regeneration in relation to 4/7/24 (Item 6 from file: 399) DIALOG(R) File 399:CA SEARCH(R) (c) 2009 American Chemical Society. All rts. reserv. 98069606 CA: 98(9)69606t JOURNAL. Serum-stimulated 3H-orotic acid incorporation into RNA by hepatocyte primary cultures AUTHOR(S): Fugassa, E.; Gallo, G.; Voci, A.; Cordone, A. LOCATION: Ist. Fisiol. Gen., Univ. Genova, 16132, Genoa, Italy JOURNAL: IRCS Med. Sci.: Libr. Compend. DATE: 1982 VOLUME: 10 NUMBER: 11 PAGES: 925-6 CODEN: IRLCDZ ISSN: 0305-6651 LANGUAGE: English SECTION: CA113006 Mammalian Biochemistry IDENTIFIERS: hepatocyte RNA formation serum stimulation, tissue culture hepatocyte serum DESCRIPTORS: Embryo, fetus... Newborn... blood serum of, RNA formation by hepatocyte in culture stimulation by Ribonucleic acid formation... by hepatocyte, in culture, blood serum stimulation of Biological transport ... of orotic acid, by hepatocyte in culture, blood serum stimulation of Animal tissue culture ... RNA formation by hepatocyte in, blood serum stimulation of Blood serum ... RNA formation by hepatocytes in culture stimulation by Liver, hepatocyte, metabolism... RNA formation by, in culture, blood serum stimulation of CAS REGISTRY NUMBERS: 65-86-1 transport of and RNA formation from, by hepatocytes in culture. blood serum stimulation of 4/7/25 (Item 7 from file: 399)

DIALOG(R)File 399:CA SEARCH(R) (c) 2009 American Chemical Society. All rts. reserv.

```
91205781 CA: 91(25)205781f
                                  JOURNAL
 Study of the effect of some purine and pyrimidine derivatives on the
 transport of RNA from hepatocyte nuclei in a cell free system
 AUTHOR(S): Zhemkova, L. N.; Nesterova, S. M.; Porollo, V. I.
 LOCATION: USSR
 JOURNAL: Farmakol. Regulyatsiya Regeneratorn. Protsessov.,
  DATE: 1979 PAGES: 327-8 CODEN: D6JOUJ LANGUAGE: Russian
CITATION:
Ref. Zh., Biol. Khim. 1979, Abstr. No. 18Ch423
 SECTION:
   CA006001 General Biochemistry
  IDENTIFIERS: RNA transport nucleus purine pyrimidine, hepatocyte
nucleus
   RNA transport
 DESCRIPTORS:
Biological transport ...
    of RNA, from hepatocyte nuclei, methyuracil and orotate effect on
Liver, hepatocyte, metabolism...
   RNA transport from nucleus of, methyluracil and orotate effect on,
   tissue proliferation in relation to
Cell nucleus...
   RNA transport from, of hepatocyte, methyluracil and orotate
effect on
Ribonucleic acids...
    transport of, from hepatocyte nuclei, methyluracil and orotate
effect
   on
 CAS REGISTRY NUMBERS:
24598-73-0 27942-00-3 RNA transport from hepatocyte nuclei response
to
 4/7/26
           (Item 8 from file: 399)
DIALOG(R)File 399:CA SEARCH(R)
(c) 2009 American Chemical Society. All rts. reserv.
  79143569 CA: 79(25)143569j
                                 JOURNAL
 Site of action of 5-fluoroorotic acid on the maturation of mouse
liver
 ribonucleic acids
 AUTHOR(S): Hadzhiolova, K. V.; Golovinski, E. V.; Hadjiolov, A. A.
 LOCATION: Inst. Biochem., Sofia, Bulg.
 JOURNAL: Biochim. Biophys. Acta DATE: 1973 VOLUME: 319 NUMBER: 3
 PAGES: 373-82 CODEN: BBACAQ LANGUAGE: English
 SECTION .
   CA906001 General Biochemistry
   CA901XXX Pharmacodynamics
 IDENTIFIERS: RNA formation inhibition fluoroorotate, ribosomal RNA
   maturation fluoroorotate
```

DESCRIPTORS:

```
Ribonucleic acids...
    formation of, by liver, fluorogrotic acid effect on
Ribonucleic acids, ribosomal...
   maturation of, by liver, fluoroorotic acid effect on
Liver, metabolism...
   ribosomal RNA maturation by, fluoroorotic acid effect on
Cell nucleus...
    5S ribosomal RNA transport from, to cytoplasm, fluocorotic acid
effect.
   on
Cytoplasm...
    5S ribosomal RNA transport to, from nucleus, fluoroorotic acid
effect
   on
 CAS REGISTRY NUMBERS:
703-95-7 ribosomal RNA maturation by liver response to
? ds
Set
       Items Description
S1
         203
               (YSBC OR OROTATE OR OROTIC) (5N) (TRANSPOR?)
          97 RD S1 (unique items)
S2
S3
          92 S2 NOT PY>2006
S4
          26
              S3 AND (GENE OR NUCLEIC OR CLONE OR POLYNUCLEIC OR
DNA)
? logoff y
       12aug09 15:15:02 User226352 Session D1162.3
           $3.84 0.621 DialUnits File5
              $12.20 5 Type(s) in Format 7
          $12.20 5 Types
    $16.04 Estimated cost File5
           $0.61
                    0.081 DialUnits File6
    $0.61 Estimated cost File6
           $0.93 0.144 DialUnits File24
               $2.70 1 Type(s) in Format 7
           $2.70 1 Types
    $3.63 Estimated cost File24
          $11.67 0.410 DialUnits File34
    $11.67 Estimated cost File34
           $0.18
                   0.023 DialUnits File40
    $0.18 Estimated cost File40
           $0.19
                   0.029 DialUnits File41
    $0.19 Estimated cost File41
           $0.39 0.076 DialUnits File45
    $0.39 Estimated cost File45
           $0.71 0.149 DialUnits File50
               $2.14 1 Type(s) in Format 7
           $2.14 1 Types
    $2.85 Estimated cost File50
           $0.41
                    0.097 DialUnits File65
    $0.41 Estimated cost File65
           $1.76 0.162 DialUnits File71
    $1.76 Estimated cost File71
```

```
$2.71 0.196 DialUnits File72
$2.71 Estimated cost File72
       $3.83 0.277 DialUnits File73
         $11.49 3 Type(s) in Format 7
      $11.49 3 Types
$15.32 Estimated cost File73
       $0.45 0.070 DialUnits File76
$0.45 Estimated cost File76
       $0.16 0.037 DialUnits File98
$0.16 Estimated cost File98
       $0.88 0.136 DialUnits File103
          $2.28 1 Type(s) in Format 7
       $2.28 1 Types
$3.16 Estimated cost File103
       $0.13 0.021 DialUnits File136
$0.13 Estimated cost File136
       $0.12 0.039 DialUnits File143
$0.12 Estimated cost File143
       $1.64 0.321 DialUnits File144
$1.64 Estimated cost File144
       $1.25 0.355 DialUnits File154
          $0.72 3 Type(s) in Format 7
       $0.72 3 Types
$1.97 Estimated cost File154
       $1.14 0.324 DialUnits File155
          $0.72 3 Type(s) in Format 7
       $0.72 3 Types
$1.86 Estimated cost File155
       $0.59 0.097 DialUnits File156
$0.59 Estimated cost File156
       $0.26
              0.055 DialUnits File162
$0.26 Estimated cost File162
       $0.54 0.039 DialUnits File172
$0.54 Estimated cost File172
       $0.26 0.018 DialUnits File305
$0.26 Estimated cost File305
       $0.08 0.021 DialUnits File369
$0.08 Estimated cost File369
       $0.11 0.031 DialUnits File370
          $1.62 1 Type(s) in Format 7
       $1.62 1 Types
$1.73 Estimated cost File370
       $0.08 0.029 DialUnits File393
$0.08 Estimated cost File393
      $14.50 1.109 DialUnits File399
         $23.84 8 Type(s) in Format 7
      $23.84 8 Types
$38.34 Estimated cost File399
       $2.23
               0.078 DialUnits File434
$2.23 Estimated cost File434
      OneSearch, 29 files, 5.042 DialUnits FileOS
$1.33 TELNET
```

\$110.69 Estimated cost this search \$110.71 Estimated total session cost 5.426 DialUnits Logoff: level 05.26.00 D 15:15:02